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Waterborne self-crosslinkable polyurethane compositions and self-crosslinkable polyurethane: acrylic hybrid compositions

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Waterborne self-crosslinkable polyurethane compositions and self-crosslinkable polyurethane : acrylic hybrid compositions

The present invention concerns a composition and a process for making selfcrosslinkable polyurethane dispersions to be used alone or as an hybrid polymer dispersion in water, said hybrid polymer dispersion being constituted by the functionalized polyurethane polymer (A) and a functionalised vinylic polymer (B) either as individual particles or as composite particles

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A isocyanate-terminated polyurethane prepolymer is first prepared in at least a high boiling point oxygenated coalescing solvent having no functional group reactive with isocyanates and that will remain after water dispersion. The composition is thereafter neutralized, dispersed in water and chain extended with a diamine.

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The polymer dispersions according to the invention exhibit a low film forming temperature associated with a very fast hardness development and a good resistance to water, solvents and stains.

The growing concerns on ecological issues have lead to the development of coating technologies having a lesser chemical impact on occupational health and environment.

Waterborne coatings constitute a very big and important category of products for the industry due to their ecological background. Intense research and development activities are ongoing in order to continue to positively impact the reduction of volatile organic compounds (VOC) like solvents and amines, but also the suppression of hazardous and ecotoxic components like some residual monomers, crosslinkers, emulsifiers, biocides, catalysts, etc.

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Polyurethane dispersions (PUD) are a fast growing class of polymers that can respond to these ecological requirements while offering a premium performance with regard to the mechanical and chemical resistance it offers on many coated substrates. The development of hybrid technologies involving the use of these polyurethane are capable to address the higher initial cost of the parent polyurethane polymer and enhance the benefits from the several polymer backbones present due to self-crosslinking properties, without the need of an external crosslinker and the disagreement of a limited pot-life before use.

The hybrid chemistries involving polyurethane dispersions have been described many times. For instance the patent US 5.541.251 describes an aqueous self-crosslinkable resin composition comprising an aqueous dispersion of at least one polyurethane polymer having anionic salt groups as sole chain-pendant functional group and at least one vinyl polymer having acetoacetoxyalkyl ester groups as lateral chains and capable of reacting with the polyurethane polymer during or after film formation, and resulting in the formation of a polymeric film having properties not found to be those of the non-functional components.

Among the most important properties requested for these waterborne dispersions are the good film formation at room temperature, and the rapid development of an optimum and non-evolutive coating hardness upon drying. This last requirement is not easily matched by the prior art.

The film formation has been a lot described, for example in "Mechanism of Film Formation from Polymer Latexes" from S.T.Eckersley, J. Coating Techn., 62, 780 (1990). It is generally admitted that the film formation happens in 3 consecutive stages: 1) water evaporation 2) capillary forces 3) particle coalescence. The particles coalescence imply the inter-diffusion of the polymer chains from two neighbour particles and was studied even in the more complicated case of the crosslinking of thermoset latex films, for example in M.A.Winnik, Journal of Coatings Technology, 74, 925, 49 (2002). A "minimum film formation temperature" (MFFT) can be determined experimentally, and constitutes an intrinsic characteristic of the polymer dispersion in water.

The drying of a waterborne coating includes the evaporation of water and the film formation from the polymer particles dispersed in water. The overall drying process of a coating can also be approached experimentally, generally by the means of slowly moving needles leaving a trace on the drying film. This method has been used a lot to study the drying of alkyds. 5 zones can be defined depending on the state of water evaporation during the film formation process: levelling > basic trace > ripped film > surface trace > dry film. This last point corresponds to the "open time" which is the time needed to reach an infinite viscosity. The drying time depends strongly on external factors (temperature, humidity, air flow, film thickness, skin formation...), but the polymer dispersion characteristics are also important (water contents, polymer nature, additives); the open time was shown to be related to the "volume-to-mass

ratio" influenced in turn by the polarity of the polymer at a given pH and by the presence of co-solvents. This phenomenon is well described in a paper called "Rheological Changes during the Drying of a Waterborne Latex Coating". F. Loflath, J.Coating Techn., 69, 867, 1997.

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Two cases can be find practically for the coating of waterborne polymers.

At elevated temperature (coatings for industrial application), the open time will be reduced to the minute- or even the second-scale, depending on the film thickness and on the oven characteristics & residence time.

At room temperature (and even down to 5°C for do-it-yourself applications), a good film formation is a prerequisite. The time scale involved for drying is around 1 hour, and this 'open time' measured with the 'needle method' is irrelevant for the application (i.e. a few minutes difference does not make a real difference to the end user).

After the immediate drying process (open time), the film is freshly formed and exhibit a certain hardness - providing in turn a relative resistance to scratch, blocking and dust pick-up - as described in "Factors Affecting Dirt Pick-up in Latex Coatings", A.Smith, J.Coating Techn., 68, 862, 1996. The time needed to reach a steady-state coating hardness at ambient temperature, and compatible with its usage is particularly relevant to the user. This property, called fast hardness, is referring to a coating which reaches rapidly an optimum and non-evolutive hardness. This is a very important requirement for do-it-yourself markets, since the user is supposed to make a normal usage of the coated substrate (f.i. wood) very rapidly after painting at room temperature (dust pick-up, blocking resistance, scratch and adhesion resistance). This phenomenon is best measured with a kinetics of pendulum hardness measurements (König, Persoz) and the time scale involved is in the range of 1hour to 7 days.

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The external parameters important to fast hardness are the temperature, the relative humidity and the coating thickness.

The state of the art does not particularly focus on the improvement of the fast hardness development although it is sometimes mentioned in some patents as one of the non-specific deliveries of the inventions. In most of the cases, the (final) hardness of the coatings from the prior art are linked to the (self)crosslinking with/without the implication of a (semi)interpenetrated network, and where air drying of the fatty acid

Empfangszeit 14.Juli 14:27 ···

compound may constitute a particular case. The balance between a good film formation (MFFT) and the rapid hardness obtained after (self)crosslinking at ambient temperature is most probably the main problem unsolved in the prior art, and that our invention aims to address with the specific usage of coalescing solvents in an original hybrid polymer composition.

The selection of solvents that remains during and after the drying of the film is a key parameter for fast hardness at ambient temperature. Indeed, it was found that the polymer nature (monomers, hardness, functionality, molecular weight, crosslinking) mainly play a role in the final hardness of the coating, rather than the hardness build-up kinetics.

The state-of-the-art process to make PUDs involves the use of N-methyl pyrrolidone (NMP) as an excellent solvent for the polyurethane prepolymer synthesis; this solvent is a small polar molecule with a high boiling point (202°C, 760 mm Hg) and has a particularly good affinity for the polymer. In the normal process, the polyurethane prepolymer in NMP is neutralized and dispersed in water and the solvent is not stripped-off from the dispersion. A further advantage of the process rely in the fact that the process solvent serves as coalescing agent for the harder polymer dispersions and provides a good film formation.

Another state-of-the-art process uses acetone or methyl ethyl ketone in the same way as before, but the solvent is then easily stripped out of the polymer dispersion due to its lower boiling point (56°C, 760mm Hg). In this case, the final dispersion is almost free of residual solvents. These dispersions provide a much better fast hardness development due to the absence of solvent. However, they might have a detrimentally higher minimum film forming temperature (MFFT) especially when applied at room temperature for do-it-yourself applications. In this case, a coalescing agent should be added to the dispersion to ensure the good film formation. Those coalescing agents are well known in the state-of-the-art, and are generally oxygenated solvent with a higher boiling point (ca 200°C, 760 mm Hg) like typically the (poly)glycol ethers and esters.

Surprisingly, it has been discovered that selected non reactive oxygenated coalescents could be used to replace the NMP in the synthesis process of the polyurethane oligomer, providing a polymer dispersion having at the same time an easy film formation and an improved fast hardness development, not obtained when the same coalescent is added to the polymer dispersion as a formulation, after the synthesis. The presence of the acetone or methyl ethyl ketone in the process is not mandatory.

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This is why the coating compositions of the present invention are based on:

- (A) a dispersion of a crosslinkable polyurethane polymer in aqueous medium, and obtainable from a polyurethane prepolymer which is the reaction product of:
- 5 (1) at least one polyisocyanate, and

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- (ii) at least one organic compound containing at least two reactive groups which can react with isocyanates, and
- (iii) at least one compound which is capable to react with an isocyanate group and which contains additional functional groups which are susceptible either to dispersion in water and/or to a crosslinking reaction
- (iv) in an oxygenated solvent (iv) selected from the coalescing agents having a higher boiling point of from 150 to 250°C, under 760 mm Hg and being chemically inert towards isocyanates during the manufacture of the polyurethane;
- the so obtained polyurethane prepolymer being further neutralised and dispersed in water, then reacted with a chain extender or capping agent (v) having or not remaining functional groups after the capping;
  - (B) a crosslinker, if the polyurethane is not self-crosslinkable
- In a prefered embodiment, the crosslinker (B) is a vinyl copolymer having functional groups reactive with functional groups of the polyurethane.

The oxygenated solvents (iv) can regroup not limitatively all those compounds responding to the basic criteria of coalescing efficiency and including water-to-polymer partitioning, evaporation rate, solvent retention, odor, color, freezing point, hydrolytic stability, biodegradability, irritancy and safety as described in the state of the art (see for instance Recent Advances in Coalescing Solvents for Water-Based Coatings, D.Randall, Polymers Paint & Color Journal, 188 (4405), 12, 14-15, 1998) The chemical inertness against the isocyanates and the ability to provide a fast hardness development upon drying and to improve of the final physico-chemical properties of the coating are both inherent to this invention.

The chemical categories of coalescing solvents (iv) responding to the above specifications include the oxygenated solvents with a balanced hydrophilic character that will tend to be preferentially sited at the particle / aqueous interface along with the surfactants, if any. They are the fully reacted alkyl or aryl esters of aromatic, aliphatic or cycloaliphatic polycarboxylic acids, the fully reacted alkyl or aryl esters of aromatic, aliphatic or cycloaliphatic polyglycols, the fully reacted alkyl or aryl ethers of

Empfansszeit 14. Juli 14:27"

aromatic, aliphatic or cycloaliphatic polyglycols, the fully reacted mixed alkyl and aryl esters and ethers of mixed aromatic, aliphatic or cycloaliphatic polyglycol-carboxylates, the neat, the alkyl and aryl substituted cyclic carbonates, the neat, the alkyl and aryl substituted cyclic esters, the neat, the alkyl and aryl substituted cyclic esters, the neat, the alkyl and aryl substituted cyclic anhydrides.

The coalescing solvents (iv) that are particularly suitable to be used in the frame of our invention are including not limitatively the dimethyl esters or dissobutyl esters of adipic, glutaric, succinic or phtalic acids and their blends, the ethyl-3-ethoxypropionate (Ektapro EEP, Eastman), the 2,2,4-trimethyl-1,3-pentanedioldiisobuttrate (Kodaflex TXBI, Eastman), the ethylene carbonate (Bp = 248°C) the propylene carbonate (Bp = 242°C), the propyleneglycol diacetate (DOWANOL PGDA, Bp = 191°C, 760 mm Hg), the dipropylene glycol dimethyl ether (PROGLYDE DMM, Bp = 175°C, 760 mm Hg). The best coalescing solvent is also non irritating (Xi-free).

The coalescing solvents (iv) can be used in an amount of 5 to 40wt%, preferably 10 to 15wt% expressed on the dry polymer, to reduce the viscosity of the prepolymer and, after dispersion in water, to ensure the good film formation and the fast hardness according to the invention.

This process may be facilitated by the addition of another optional low boiling point solvent, in order to further reduce the viscosity if this would appear to be necessary. Suitable solvents, either alone or in combination, are those belonging to ketones with a relatively low boiling point so that they can easily be removed before, during or after the chain extension by distillation under reduced pressure. Examples of such solvents include acetone, methyl ethyl ketone, diisopropyl ketone, methyl isobutyl ketone.

Advantages of the invention is the extremely good fast hardness development of the coating, associated with a low MFFT and a good performance profile (stain resistance, solvent resistance, etc)

Another advantage is the reduced level of the coalescing solvent necessary to get a workable viscosity suitable for dispersion in water.

The absence of any stripping operation of a process solvent is also an advantage, with a positive impact on productivity.

Finally, the use of a coalescing solvent is not necessarily irritating when compared with typical solvents used for making polyurethanes, such as N-methyl pyrrolidone.

Empfangszeit 14. Juli 14:27

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The preparation of the polyurethane prepolymer bearing terminal isocyanate moieties can be carried out in a conventional manner, by reacting a stoichiometric excess of the organic polyisocyanate(s) with all the organic compounds containing at least two reactive groups which are enabled to react with isocyanate groups under substantially anhydrous conditions, preferably at a temperature between 50°C and 120°C., more preferably between 70°C and 95°C., until the reaction is complete. The reaction may be carried out in the presence of any of the known catalysts suitable for polyurethane preparation such as amines and organometallic compounds. Examples of these catalysts include triethylenediamine, N-ethyl-morpholine, triethylamine, dibutyltin dilaurate, stannous octanoate, dioctyltin diacetate, lead octanoate, stannous oleate, dibutyltin oxide and the like.

During the preparation of the isocyanate-terminated polyurethane prepolymer the reactants are generally used in proportions corresponding to an equivalent ratio of isocyanate groups to such groups which are enabled to react with the isocyanate functionalities of from about 1.1:1 to about 2:1, preferably from about 1.6:1 to 1.8:1. The polyurethane prepolymer thus contains terminal free isocyanate groups, because the polyisocyanate is used in excess and the polyurethane polymer can be obtained from the polyurethane prepolymer by neutralization and dispersion in water, followed by the further reaction with a capping agent such as water or a chain extender.

An important feature of the aqueous compositions of the present invention is that the compositions are crosslinkable and that during application or preferably after application of the composition to a substrate, a three-dimensional network can be formed by crosslinking the composition.

In the basic embodiment, the composition is rendered crosslinkable by incorporating reactive groups into the polyurethane polymer which are capable of reacting either with each other or with a crosslinking agent which preferably is also a compound of the aqueous coating composition.

In another embodiment, the polyure hance polymer is obtained from the reaction of a polyurethane prepolymer with a capping agent which contains an additional functionality which is susceptible to a thermal or radiation initiated crosslinking reaction.

The dispersion may also optionally contain a thermal initiator or photo-initiator for radical or cationic polymerization.

Empfansszeit 14. Juli 14:27

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In still another embodiment, the polygrethane does not contain a special functionality which is susceptible to a self-crosslinking reaction, but the aqueous coating composition contains a crosslinking agent which can react with the polygrethane polymer to form a three-dimensional molecule.

In the most preferred embodiment, the dispersion in water also contains a crosslinking agent which preferably is a reactive functionalized oligomer or polymer other than the polymerthane polymer.

The polyisocyanate (compound i) used according to the present invention for the preparation of the polyurethane prepolymer may be an aliphatic, cycloaliphatic, aromatic or heterocyclic polyisocyanate or a combination thereof. As example for suitable aliphatic diisocyanates there may be mentioned 1.4-diisocyanatobutane, 1.6-dilsocyanatohexane, 1.6-dilsocyanato-2.2.4-trimethylhexane, and 1.12-diisocyanatododecane either alone or in combination. Particularly suitable cycloaliphatic diisocyanates include 1:3- and 1.4-diisocyanatocyclohexane, 2.4-diisocyanato-1-methyl-cyclohexane, 1.3-diisocyanato-2-methylcyclohexane, 1-isocyanato-2-(isocyanatometyl)-cyclopentaine, 1.1'-methylenbis[4-

isocyanato-2-(isocyanatometyl)-cyclopentable, 1,1'-(1-methylefhylidene)bis[4-isocyanato-cyclohexane], 5isocyanatocyclohexanel. isocyanato-1-isocyanatomethyl-1,3,3 trimethylcyclohexane (isophorone diisocyanate), 1,3- and 1,4-bis(isocyanatomethy))cyclomexane, 1,1'-methylene-bis[4-isocyanato-3-1-isocyanate-4(or 3)-isocyanatomethyl-1-methylcyclohexane methylcyclohexane], either alone or in combination. Particularly suitable aromatic diisocyanates comprise 1.4-diisocyanatobenzene, 1.1'-metilylenebis[4-isocyanatobenzene], 2.4-diisocyanato-1-mthylethylidene)bis[4-isocyanatolicnzene], 1,3and 1.4-bis[1-isocvanato-1methylethyl)benzene, 1,5-naphtalene ditsocyanate, either alone or in combination. Aromatic polyisocyanates containing 3 of more isocyanate groups may also be used such as 1,1',1"-methylidynetris[4-isocyanatobenzene] and polyphenyl polymethylene polyisocyanates obtained by phosgenation of aniline/formaldehyde condensates.

The total amount of the organic polyisogranate is not particularly restricted, but generally is in the range from 10 to 60 wt% of the polyurethane polymer, preferably from 20 to 50 wt% and more preferably from 30 to 40 wt%.

In a preferred embodiment said polyisocyanate is selected from cycloaliphatic polyisocyanates, especially preferred is the use of methylene-bis(cyclohexyl isocyanate).

Empfangszeit 14.Juli 14:27

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The organic compounds containing at least two reactive groups which can react with isocyanates (compound ii) are preferably polyols, but e.g. amines can also be used.

Suitable examples are polyester polyols, polyether polyols, polyacetal polyols, polyesteramide polyols, polyacrylate polyols, polythioether polyols and combinations thereof. Preferred are the polyester polyols, polyether polyols and polycarbonate polyols. These organic compounds containing at least two reactive groups which are enabled to react with isocyanates, preferably have a number average molecular weight within the range of 400 to 5,000, preferably 600 to 1,000.

Polyester polyols are particularly preferred and suitable polyester polyols which may be used comprise the hydroxyl-terminated reaction products of polyhydric, preferably dihydric alcohols (to which trihydric alcohols may be added) with polycarboxylic, preferably dicarboxylic acids or their corresponding carboxylic acid anhydrides. Polyester polyols obtained by the ring opening polymerization of lactones may also be used. The polyester polyol may also contain an air-drying component such as a long chain (un)saturated fatty acid.

The polycarboxylic acids which may be used for the formation of these polyester polyols may be aliphatic, cycloaliphatic, aromatic and/or heterocyclic and they may be substituted (e.g. by halogen atoms) and saturated or unsaturated. As examples of aliphatic dicarboxylic acids, there may be mentioned, succinic acid, glutaric acid, adipic acid, suberic acid, azelaic acid sebacic acid and dodecanedicarboxylic acid. As an example of a cycloaliphatic dicarboxylic acid, there may be mentioned hexahydrophthalic acid. Examples of aromatic dicarboxylic acids include isophthalic acid, terephthalic acid, ortho-phthalic acid, tetrachlorophthalic acids and 1,5-naphthalenedicarboxylic acid. Among the unsaturated aliphatic dicarboxylic acids which may be used, there may be mentioned fumaric acid, maleic acid, itaconic acid, citraconic acid, mesaconic acid and tetracarboxylic acids include trimellitic acid, trimesic acid and pyromellitic acid.

The polyhydric alcohols which are preferably used for the preparation of the polyester polyols include ethylene glycol, propelene glycol, 1,3-propanediol, 1,3-butanediol, 1,4-butanediol, 1,5-pentanediol, 1,6-hexanediol, neopentyl glycol, diethylene glycol, dipropylene glycol, triethylene glycol, tetraethylene glycol, dibutylene glycol, 2-methyl-1,3-pentanediol, 2,2,4-trimethyl-1,3-pentanediol, 1,4-cyclohexanedimethanol, ethylene oxide adducts or propylene oxide adducts of bisphenol A or hydrogenated

Empfangszeit 14.Juli 14:27

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bisphenol A. Triols or tetraols such as trimethylolethane, trimethylolpropane, glycerin and pentaerythritol may also be used. These polyhydric alcohols are generally used to prepare the polyester polyols by polycondensation with the above-mentioned polycarboxylic acids, but according to a particular embodiment they can also be added as such to the polyurethane prepolymer reaction mixture.

In a preferred embodiment the polyester polyol is made from the polycondensation of neopentylglycol and adipic acid.

Suitable polyether polyols comprise polyethylene glycols, polypropylene glycols and polytetramethylene glycols, or bloc copolymers thereof.

Suitable polycarbonate polyols which may be used include the reaction products of diols such as 1.3-propanediol. 1.4-butanediol, 1.6-hexanediol, diethylene glycol, triethylene glycol or tetracthylene glycol with phosgene, with dialkylcarbonates or diarylcarbonates such as dimethylcarbonate or diphenylcarbonate or with cyclic carbonates such as ethylene and/or propylene carbonate.

Suitable polyacetal polyols which may be used include those prepared by reacting glycols such as diethyleneglycol with formaldehyde. Suitable polyacetals may also be prepared by polymerizing cyclic access.

The total amount of these organic compounds containing at least two reactive groups which can react with isocyanates preferably ranges from 30 to 90wt% of the polyurethane polymer, more preferably of from 45 to 65wt%.

The at least one compound which is capable to react with an isocyanate group and which contains additional functional groups (compound iii) is an alcohol or a polyol having a pendant functionality. Sich an alcohol or polyol typically contains water soluble side chains of ionic or con-lowic nature suitable to allow the polymer dispersion in water. Preferably, the polyol has functional groups such as anionic salt groups or similar precursors which may be subsequently converted to such anionic salt groups, such as carboxylic or sulfonce acid groups. It is also possible that the polyol comprises other functional groups which are susceptible to a crosslinking reaction, such as isocyanate, hydraxy, amine, acrylic, allylic, vinyl, alkenyl, alkinyl, halogen, epoxy, aziridine, aldesyde, ketone, anhydride, carbonate, silanol, acetoacetoxy, carbodilmide, ureidoa kyl, hamethylolamine, N-methylolamide N-alkoxymethyl-amine, N-alkoxymethyl-amine, or the like.

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Compounds which are capable of reacting with isocyanate groups and containing anionic salt groups (or acid groups which may be subsequently converted to such anionic salt groups) preferably are the compounds containing the dispersing anionic groups which are necessary to render the polyurethane prepolymer self dispersible in water e.g. sulfonate salt or carboxylate salt groups. According to the invention, these compounds are preferably used as reactions for the preparation of the isocyanate-terminated polyurethane prepolymer.

The carboxylate salt groups incorporated into the isocyanate-terminated polyurethane . 10 prepolymers generally are derived from polyhydroxycarboxylic acids represented by the general formula (HO)xR(COOH)y, wigerein R represents a straight or branched hydrocarbon residue having 1 to 2 camon atoms, and x and y independently are integers from 1 to 3. Examples of these hydroxycarboxylic acids include citric acid and tartaric acid. The most preferred hydroxycarboxylic acids are the a,a-15 dimethylolalkanoic acids, wherein 2 and y in the above general formula, such as for example, the 2,2-dimethylolpropionic acid. The pendant anionic salt group content of the polyurethane polymer may virilia wide limits but should be sufficient to provide the polyurethane with the e reguired degree of water-dispersability and crosslinkability (if no other crosslinkabile group is incorporated in the polyurethane 20 polymer which provides the required crosslinkability). Typically, the total amount of these anionic salt group-containing compounds in the polyurethane polymer can range from 1 to 25wt% of the polymethane polymer, preferably from 4 to 10wt%.

The sulfonate salt groups can be introduced in this prepolymer using sulfonated polyesters obtained by the reaction of sulfonated dicarboxylic acids with one or more of the above-mentioned polyhydric alcohols, on by the reaction of sulfonated diols with one or more of the above-mentioned polycarboxylic acids. Suitable examples of sulfonated dicarboxylic acids include 5-(sodiosulfo)-isophthalic acid. Suitable examples of sulfonated diols include sociosilfonydroquinone and 2-(sodiosulfo)-1,4-butanediol.

Any acid functionality which may be present in the polyurethane prepolymer can be converted to anionic salt groups by neutralization of said groups, before or simultaneously with the preparation of an aquieous dispersion of this prepolymer. The dispersion process of the polyurethane prepolymer is well known to those skilled in the art, and usually requires rapid mixing with a high shear rate type mixing head. Preferably, the warm or cold polygrethane prepolymer is added to the water under

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vigorous agitation or, alternatively water may be stirred into the warm or cold polyurethane prepolymer.

Suitable neutralizing or quaternizing agents for converting the above mentioned acid groups of the polyurethane prepolymers into anionic salt groups during or before the dispersion in water can be volatile organic bases and/or non-volatile bases, especially triethylamine.

Suitable volatile organic bases cambe preferanly selected from the group comprising ammonia, trimethylamine, triethylamine, triisopropylamine, tributylamine, N.N-dimethylcyclohexylamine, N.N-dimethylamine, N-methylmorpholine, N-methylpiperazine, N-methylpyrrolidine and N-methylpiperidine. The trialkylamines are preferred.

Suitable non-volatile bases include those comprising monovalent or bivalent metals, preferably alkali metals such as likihium, sodium, potassium or calcium. These nonvolatile bases may be used in the form of inorganic or organic salts, preferably inorganic salts wherein the anions do not remain in the dispersions such as metal hydroxides, carbonates and bicarbonates.

The total amount of these neutralizing agents should be stoechiometrically calculated according to the total amount of accingroups to be neutralized. To ensure that all acid groups are neutralized in the case volatile organic bases are used, it is advisable to add the neutralizing agent in an excess of p to 30wt%, preferably 0 to 10wt%.

The aqueous polyurethane polymer is prepared by dispersing the neutralized polyurethane polymer in water or are massively the addition of water to the neutralized prepolymer solution, until phase inversion occurs and water becomes the dispersing phase.

If the functional groups are acidic groups which should be transformed to anionic groups, it can be preferable that the neutralizing reaction of the acidic groups is performed before the polyurethane preportation is dispersed into the aqueous medium. However, it is also possible that the aqueous medium into which the polyurethane polymer is dispersed contains the neutralizing agent.

Polyurethane prepolymer dispersions after chain extension of the free isocyanate end

Empfansszeit 14.Juli 14:27

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groups with a capping agent (v), wherein the capping agent is a well known agent used to inactivate the terminal isocyana groups. The capping agent can e.g. be water or a usual chain extender.

The chain extender should carry active highrogen atoms which react with the terminal 5 isocyanate groups of the polyuretiane propolymer. The chain extender is suitably a water-soluble aliphatic, alicyclic, aromatic or heterocyclic primary or secondary polyamine having up to 80, preferally up 12 carbon atoms.

When the chain extension of the posturethane prepolymer is effected with a 10 polyamine, the total amount of polyamine should be calculated according to the stoechiometricamount of isocyanategroups present in the polyurethane prepolymer in order to obtain a fully reacted polymerthane polymer (a polymerthane urea) with no residual free isocyanate groups; the polyamine used in this case may have an average 15 functionality of 2 to 4, preferably 2. 3,

The degree of non-linearity of the polymer is controlled by the functionality of the polyamine used for the chain extension. The desired functionality can be achieved by mixing polyamines with different amine functionalities. For example, a functionality of 2.5 many behachieved by using equimolar mixtures of diamines and triamines.

Examples of such chain extenders useful herein comprise hydrazine, ethylene diamine, piperazine, diethylene priamine, triethylene tetramine, tetraethylene 25 pentamine, pentaethylene piperazinoethyl)ethylenediamine, aminoethyl)ethylenediamine, aminoethyl)piperazine. N-(2-amino bis(2-aminoethyl)-N-(2-piperazinoetsyl)amine. guanidine, melamine, N-(2-aminoginyi)-13-propanediamine, 3,3'-diaminobenzidine 30 2,4,6-triaminopyrimidine, dig opyie etriamine. tripropylenetetramine, N.N-bis(6-aminoheayi)amine, N,N'-bis(3-aminopropyl)ethylenediamine. 2.4-bis(4'-aminobenzyl)amine, in-butanediamine, 1,6-hexanediamine, 1,8octanediamine, dodecanediamine, isophorone dia bls(4-aminocyclohexyl)memane cyclohexane), bis(4-amino-3 methacyclohexyl)methane methanel. methylcyclohexane-4-yl)methane].

N,N,N-tris(2-aminoethyl)amine. N-(2-W.N-05(2-aminoethyl)piperazine, N,N,N'-tris(2-N-[N-(2-aminoethyl)-2-aminoethyl]-N'-(2hyl)- - (2-piperazinoethyl)ethylenediamine, N.N. N.N-bis(2-piperazinoethyl)amine, tetrapropylenepentamine, 1,10-decanediamine, 2-methylpentamethylenediamine, ine pr 1-amino3-aminomethyl-3,5,5-trimethyl-[OT bis(aminocyclohexane-4-y1)-[or bis(amino-2alpha. omega-polypropyleneglycol-diamine-

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sulfopropylated sodium salts, polyahyleze amines, polyoxyethylene amines and/or polyoxypropylene amines (e.g. Jeffaggines grom TEXACO). In a preserred embodiment the chain extender is selected from aliphatic diamines.

preferably it is 1,5-diamino-2-methyl pengine.

The total amount of polyamines should be calculated according to the amount of isocyanate groups present in the polyula hane prepolymer. The ratio of isocyanate

groups in the prepolymer to active aydresen in the chain extender during the chain extension may be in the range of from all it 1.0:0.7 to about 1.0:1.1, preferably from about 1.0:0.9 to about 1.0:1.02 on an equivalent basis.

The chain extension reaction is generally carried out at a temperature between 5° and 90°C, preferably between 10° to 50 c. and most preferably between 15° to 20° C.

15 20 susceptible to a crosslinking reacti may in addition also be used for game compounds are aminopropyltriethoxysilane, the Nipeta-alignoethyl-gamma-25

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In another embodiment of the present in mation, the chain capping agent contains the reactive groups which are capable of electing the crosslinking of the polyurethane polymer during or after application of the aqueous coating composition to the substrate. In this case, it is possible that the prepolymer is prepared by only three components and does not contain the at least one compound which is capable to react with an isocyanate group and which contains additional functional groups which are (compound iv), but, of course, such a compound pregaring the prepolymer. Examples of these -am propyltrimethoxysilane. gamma-

aminopropyltrimethoxysilane, the bis-(gamma-trimethoxysilylpropyl) amine, the Nbeta-(aminoethyl)-gamma-aminopippylmathyldimethoxysilane, or the like.

polymerisation

diamines as chain extenders. polypropylenegiycoi-diamine.

In still another embodiment, the chain apping agent can be a poly(meth)acrylate compound capable to crossling after thermal or radiation induced radical

In a further preferred embodiment of the present invention, sulfonates groups can be incorporated into the polyurethane polymer by a chain extension using sulfonated mple the sodium salt of 2,4-diamino-5ex methylbenzenesulfonic acid or the sodium salt of sulfopropylated alpha, omega-

The aqueous coating compositioning the present invention can be obtained from the sole polyurethane dispersion desembed prove. Nevertheless, it may also contain at

Empfangszeit 14.Juli 14:2

least one external crosslinking age compounds which can react with the melamine-formaldehyde resin, as described in US-A 4,598,121, to which it is referred for details.

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toxic and have negative effects on functional and reactive vinyl-ty

typically reserved for vinyl-type poly decrease of the minimum film forms radical addition polymerization of aninyl pe monomer.

polymerisation technique, prefera miniemulsion polymerisation.

100,000 and 500,000.

methods, for example by using a

at, especially if the functionality present on the polymer is not sufficient to provide self-crisslinking. The term "crosslinking agent" as used in the present specification is no restrictive and encompasses all kinds of polymer, preferably with functional groups of the polyurethane polymer to a three-dimensional network. Suitable crosslinking agents are known in the polyurethane contains carboxyl groups as functional groups which are susceptible to a crosslinking reaction, the crosslinking agent can be a trifunctional aziridine compound or a t is described in US-A 4,301,053 and US-A 5,137,967, to which it is referred for detail. If the additional functional groups which are susceptible to a crosslinking agaction are obtained by incorporating hydrazide groups into the polyurethane change the prosslinking agent can be formaldehyde, as

Since crosslinking agents such as aziridize compounds or formaldehyde are relatively e position, it is preferred to use e privmers as crosslinking agents for the polyurethane dispersion. They have the agaitional benefit to (i) add additional features iners like an excellent aging (ii) participate to the decrease of the minimum film formation emperature without sacrifying the hardness of the polyurethane component or wairing and amounts of coalescing solvents (iii) offer an improved performance due to the crosslinking reaction without generating restrictions linked to a limited possife of to toxicity problems. The term "vinyl-type" polymer as used in the present specification is not specifically restricted and should encompass all types of polymers btanzble by polymerization, preferably by free

The vinyl-type polymer may be prepared by any suitably free-radical initiated by mulsion polymerization or alternatively by

The vinyl-type polymers for use in the present invention may preferably have a weight average molecular weight within the range of 10,000 to 500,000, preferably between range of 10,000 to 500,000, preferably between

The emulsion polymerisation of the morn may be carried out according to known mi-lanch process wherein a pre-emulsion of the above-mentioned monomers is ingroduced into a reactor containing an aqueous

Empfangszeit 14.Juli 14:27

solution of a free-radical initiator and heated at a constant temperature between 60° and 95°C, preferably between 75° and 87°C, for a period of 1 to 4, preferably 2 to 3 hours to complete the reaction.

The pre-emulsion of the monomers can be prepared by adding each monomer with stirring to an aqueous solution of an emit sifier, preferably an anionic type emulsifier, such as for example lauryl sulfate dode cylbenzenesulfonate, dodecyl diphenyloxide disulfonate, alkylphenoxypoly(ethylenexy)- or (propyleneoxy)sulfates and combination thereof or dialkylsulfosucce ates, wherein the alkyl residue may have from 8 to 12 carbon atoms. Most preferably, a nonylphenoxypoly(ethyleneoxy)sulfate is used. It is to be understood that non-io ic emulsifiers may also be used.

Conventional free-radical initiators are used for the polymerisation of the monomers, such as for example hydrogen perox de, tert-butylhydroperoxide. alkali metal persulfates or ammonium persulfate

Vinyl-type monomers are generally ethylenically unsaturated, preferably monoethylenically unsaturated monomers. Preferred ethylenically unsaturated monomers which may be used for the formation of the vinyl-type polymer are selected from the group comprising

- a) o.ß-monoethylenically unsaturate: carboxylic acid and their esters like alkyl acrylates and alkyl methacrylates, which have an alkyl residue of 1 to 12 carbon atoms, such as methyl methacrylate methyl acrylate, ethyl acrylate, butyl acrylate, 2-ethylhexyl acrylate, isooctyl acrylate nony, acrylate and dodecyl acrylate,
- b) a \$\beta\$-monoethylenically unsaturate carboxylic acid and their functionalised esters like hydroxyalkyl acrylates and hydroxyalkyl methacrylates, which have an alkyl residue of 1 to 12 carbon atoms, such as hydroxyethyl acrylate, hydroxyethyl methacrylate.
- c) vinyl substituted aromatic hydroca bons such as styrene, a-methylstyrene and the like.
- 35 d) a.B-ethylenically unsaturated trarbonamides such as acrylamide, methacrylamide, methoxymethylacrylamide N-methylolacrylamide and the like.

Empfansszeit 14.Juli 14:27

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- e) vinyl esters of aliphatic acids such as vinyl acetate, vinyl versatate and the like (versatates are esters of tertiary monocar explic acids having C9, C10 and C11 chain length).
- 5 f) · vinyl chloride and vinylidene chloride,

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g) monoethylenically unsaturated sufforates such as the alkali metal salts of styrenesulfonic acid, 2-acrylamido-2-methyl-propanesulfonic acid, 2-sulfoethyl methacrylate, 3-sulfopropyl methacrylate and the like (internal surfactants).

Necessarily, at least one of said monomers must contain a functional group chosen between carboxylic and sulfonic acids. In particular, allylic, vinyl. alkenyl, alkinyl, halogen, carbonate, silane, acetoacetoxy. Carbonate, silane, aldehyde, ketone, anhydride, carbonate, silane, acetoacetoxy. Carbonate, silane, aldehyde, ketone, anhydride, carbonate, silane, acetoacetoxy. Carbonate, silane, aldehyde, ketone, anhydride, carbonate, silane, acetoacetoxy. Carbonates, hydroxy, amine, acrylic, allylic, epoxy aziridine, aldehyde, ketone, anhydride, carbonate, silane, acetoacetoxy. Carbonates, hydroxy, amine, acrylic, allylic, epoxy aziridine, aldehyde, ketone, anhydride, carbonate, silane, acetoacetoxy. Carbonates, hydroxy, amine, acrylic, allylic, epoxy aziridine, aldehyde, ketone, anhydride, carbonates, hydroxy, amine, acrylic, allylic, epoxy aziridine, aldehyde, ketone, anhydride, carbonates, hydroxy, amine, acrylic, allylic, epoxy aziridine, aldehyde, ketone, anhydride, carbonates, hydroxy, amine, acrylic, allylic, epoxy aziridine, aldehyde, ketone, anhydride, carbonates, hydroxy, amine, acrylic, allylic, epoxy aziridine, aldehyde, ketone, anhydride, carbonates, hydroxy, amine, acrylic, allylic, epoxy aziridine, aldehyde, ketone, anhydride, carbonates, hydroxy, amine, acrylic, allylic, epoxy aziridine, aldehyde, ketone, anhydride, carbonates, hydroxy, amine, acrylic, allylic, epoxy aziridine, aldehyde, ketone, anhydride, carbonates, hydroxy, amine, acrylic, allylic, epoxy aziridine, aldehyde, ketone, anhydride, carbonates, hydroxy, amine, acrylic, allylic, epoxy aziridine, aldehyde, ketone, anhydride, carbonates, hydroxy, amine, acrylic, allylic, epoxy aziridine, aldehyde, ketone, anhydride, carbonates, hydroxy anhydrates, hydroxy axiridine, aldehyde, keto

It can be advantageous to use an emulsion polymerization process with a sequential monomer addition feed known to those skilled in the art in order to obtain core & shell or assimilated particle morphology that help the film formation process while giving at the same time an improved hardness and resistance of the coating.

In a preferred embodiment of the present swention, the aqueous coating composition contains a crosslinking agent, wherein the crosslinking agent is a vinyl-type monomer as described above which contains as functional groups chain-pendant acetoacetoxyalkyl ester groups capable to react with the polyurethane polymer containing anionic salt groups, preferable carboxylate salt groups. Such vinyl-type polymers with chain-pendant acetoacetox alkyl ester groups can e.g. be prepared by polymerizing vinyl-type monomers as discussed above, wherein at least a part of the vinyl-type monomers contains an aceto-acetoxyalkyl ester group. In a preferred embodiment, the vinyl-type monomers have the general formula R1-O-CO-CH2-CO-CH3 wherein R1 represents a CH2=CR'-Cl D-R"-group or a CH2=CR'R"-group in which

. Empfangszeit 14.Juli 14:27

R' is -H or -CH3, and R" is an alkyl residue having 1 to 12 carbon atoms. The most preferred monomers of this type are acete acetoxyethyl acrylate and acetoacetoxyethyl methacrylate.

- The amount of the monoethylenically unsaturated monomer containing an acetoacetoxyalkyl ester group may generally vary from about 1 to about 80wt%, preferably from about 5 to 50wt% and most preferably 5 to 20wt% of the vinyl polymer.
- Thus, the preferred crosslinking agent is a vinyl-type polymer comprising chainpendant acetoacetoxyalkyl ester functional groups, preferably formed by the freeradical addition polymerisation of at least one monoethylenically unsaturated
  monomer containing an acetoacetoxyalkyl ester group with at least one other
  ethylenically unsaturated monomer as defined above.

Vinyl-type polymers containing chain pendant functional acetoacetoxyalkyl ester groups and methods for producing such polymers are e.g. disclosed in US-A 5,541,251.

The vinyl-type polymer can be combined with the polyurethane polymer in an aqueous composition by dispersing both compounts in an aqueous medium, preferably water.

In another embodiment, the vinyl-type polymer is formed in situ by polymerizing one or preferentially several vinyl-type monomers in the presence of an aqueous polyurethane dispersion. Alternatively, it salso possible to prepare the polyurethane polymer in the presence of the vinyl-type polymer, by subjecting an isocyanate-terminated polyurethane prepolymer having anionic salt functional groups to chain-extension with a capping agent in the presence of an aqueous dispersion of a vinyl polymer having functional groups.

Finally, it is possible to mix the precipiner with vinyl type monomers and to polymerise in the solvent phase (or even ally to add a preformed vinyl type polymer in the solvent phase and then to neutralise and disperse in water the intimate polymer mix into hybrid particles.

Thus, in the most preferred embodiment of the present invention, the polyurethane polymer contains additional functional groups which are susceptible to a crosslinking reaction and which are an anionic salit group, preferably a group COOM or SO3M,

Empfangszeit 14. Juli 14:27

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wherein M represents an alkali metallor an ammoniumtetraalkylammonium or tetraalkylphosphonium group, as define require expensive and potentially toxic crisslinking agents.

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In an embodiment of the present invertion described above, the aqueous coating composition comprises the polyurethan polymer and the vinyl-type polymer in a

The aqueous coating composition of the gresent invention can comprise additional functionality in the polyurethane polymen by a skilled person accordingly.

If desired, the compositions of the present invention may include other auxiliary 25 include biocides, antioxidants, plasticize foam control agents, thickening agents, 30 polypropylene, polystyre polyvinylidene chloride, polyacryjate dispersions. 35

The crosslinking agent and optional auxiliary substances or additives are included into the aqueous dispersion by stirring in a known manner.

in US-A 5,541,251 and the crosslinking agent is a vinyl-type polymer having chall pendant acetoacetoxyalkyl ester functional groups, whereby crosslinking is effected at moderate temperatures during and/or after film-formation. These compositions have a remarkably long pot-life and do not

weight ration of 10:0 to 1:10, preferably one 4:1 and more preferably of 1:2 to 2:1.

crosslinking agents, e.g. polyfunctional pholecules having reactive functionalities including carboxylic and sulfonic acids, secvanates, hydroxy, amine, acrylic, allylic, vinyl, alkenyl, alkinyl, halogen, epoxyl aziridine, aldehyde, ketone, anhydride, carbonate, silane, acetoacetoxy, carbo limide, ureidoalkyl, N-methylolamine, N-methylolamide N-alkoxy-methyl-amine, light and like these other crosslinking agents may be presentill the aqueous coating composition alone or in combination with one another or with the vinyl-type polymer as discussed above. Which crosslinking agent should be used depends on the type of crosslinkable and the crosslinking agent can be chosen

substances (additives) which may be added to the final composition in order to impart or improve desirable properties or to suppliess undesirable properties. These additives colorants, pigments, silica sols and the known surfactants, leveling agents, wetting agents, humectants, antifoaming agents, Palescing agents, heat stabilizers, UV-light stabilizers, waxes, etc. The composition have also be blended with other polymer dispersions in water, for example proving acetate/versatate, epoxy resins, polybutadiene, polyvinyl other homopolymer and copolymer

The aqueous coating compositions suitabsenave a total solids content of from about 5 to 65wt%, preferably from about 30 to 5 wt% more preferably from 30 to 35wt%; a viscosity measured at 25°C of 10 to 5000 mpa s, preferably 100 to 500 mpa s, a pH value of 7 to 11, preferably of 7 to 9 and 22 average particle size of about 10 to 1000 nm, preferably 80 to 300 nm, more preferably 50 to 150 nm. The film formation temperature may preferably range from 0 to 25°C, more preferably from 0 to 5°C.

The adhesive or protective aqueous coating composition of the invention can be easily applied to wood, which is the primary application for this invention, but also to any other substrate including not limitatively paper, cardboard, plastics, fabrics, glass, glass fibers, ceramics, concrete, leather, in stals and the like, for industrial or domestic purposes and by any conventional method including brushing and roll coating, but also spraying, dipping, flexography are thelrography, at room temperature or at elevated temperature.

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The following examples are given for illustrating the invention.

The dry content was measured by a greatment method and expressed in %. The drying procedure requests 2 hours at 16 c.

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The pH is measured using a conventional at intermeter fitted with a glass electrode.

The viscosity (n) of the aqueous polymer dispersions was measured at 25°C with a Brookfield RVT Viscometer, using spinds No. 1 at 50 rpm when the viscosity was under 200 mPa s or spindle No. 2 at 50 mm when the viscosity was higher than 200 mPa s, and is expressed in mPa.s

The average particle size of the aqueous folymer dispersions was measured by laser light scattering using a Malvern Perticle analyzer Processor types 7027 & 4600SM, and is expressed in nm.

The grits value is the amount of residue from the polymer dispersion filtered on a  $50\mu$  sieve and is expressed in mg/liter.

The polymer blends are prepared under socient agitation and formulated accordingly with UCECOAT XE 430 (thickener) in one to obtain a viscosity of about 1000 mPa.s and with a coalescing agent only when specified in the text.

Empfangszeit 14. Juli 14:27

Except when otherwise stated, the waters sed formulations are applied on wood using a brush: 3 consecutive uniform layers see prepared with a drying time of 1 hour between the layers always followed by a sattle rubbing with sand paper. After having been applied to the substrate, the 3-layers decatings are allowed to cure at ambient temperature (23°, 50% relative humidity) as a period of 7 days.

The König hardness is assessed according to DIN 53157 as follows: a 100  $\mu$  (wet) coating is applied on a glass plate using a Meyer bar. The film is placed in a conditioned room (23°C, 50% humidity) and is allowed to dry. A measurement of the hardness is made after 1hour, 4hours, 1 sy. 4 days, 7 days using a König pendulum equipment. The pendulum records the scillation time in seconds (s) between two fixed angles. High oscillation values correspond to high coatings hardness.

The Minimal Film Formation Temperature (MFFT) is assessed as follows: a wet coating is applied on a stainless steel plate with a stable temperature gradient and covered with a top cover to prevent air circula on. The film is allowed to dry, and the minimum temperature delivering a uniform and scratch resistant coating is recorded.

The quality of the film is assessed as follows: a coating is made on wood using a brush and allowed to dry at 23°C and 50% hursaity. After 24 hours, the coating is dry and the quality of the film formation is observed using a microscope in order to detect micro-crack defects. The same test can be performed at 5°C. A quotation is given on a 1-5 scale (5 = no defects).

The clarity is assessed as follows: a 100 evet) coating is applied on a black reference substrate (Leneta) and on a glass plate using a Meyer bar. The film is placed in a conditioned room (23°C, 50% humidity) evel is allowed to dry. A visual assessment of the film haze and/or whiteness (on Leneta) and the film transparency (on glass) is made and a quotation is given on a 1-5 state (1 = white, hazy 5 = transparent). A high clarity is expected to provide coatings with an excellent and decorative aspect.

The stain resistance of a coating is lasses to by putting a test substance covered with a microscope glass on the coating and let for a hours (1 hour for hot water). The test substances used are cold and hot water mustard, ammonia, ethanol and red wine. The stains are washed after a couple of rubs using a tissue saturated with isopropanol. The remaining stains are as seed visually using a 1-5 scale, 5 = best. A high value (5) is expected to provide the best protection against any household product spillage.

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The solvent resistance is assessed by douter rubs made by pressing the coating with a . peace of cotton rag saturated with water manol 1:1 or with acetone while applying a forward and backward motion. The reported humber is the number of double rubs required to break through the coating. A the value (>100) is expected for an optimum resistance.

The scratch resistance of the coatings is a sessed using a coin which is firmly pressed and moved over the coating. The result the damage observed is referred to a 1-5 scale, 5 = best. A high scratch resistance negessary to impart long term mechanical protection to the substrate

The adhesion was measured using an ad sive tape firmly pressed on a cross-cutted coating area and removed rapidly; the state wet test can be made when a drop of water is in contact with the cross-cutted sating area for 2 hours. The damage to the coating due to adhesion loss is expressed a a 5-5 scale, 5 = best. A high adhesion (5) is necessary to ensure a strong permerent bond between the coating and the substrate.

The isocyanate content in a polyurethane repolymer reaction mixture was measured 20 using the dibutylamine back-titration met and is expressed in meq/g.

In examples 1-3, only the process and solvent usage are modified (i.e. the polyurethane polymer composition remains junchanged) In examples 4-8, the composition of the polymers is being varie

Comparative example. Reference synthesis of Pul 1 with standard process using NMP.

A double-wall glass reactor equipped with a mechanical stirrer, a thermocouple, a 30 vapor condenser and a dropping fullel was charged with 216.2 g of Nmethylpyrrolidone. 141.5 g of a polyeste diol having an average molecular weight -670 Daltons and obtained by the polycondensation of adipic acid and neopentylglycol, 42.6 g of cyclohexane dir hamol, 41.0 g of dimethylol propionic acid. 383.9 g of methylene bis(cyclohexyl isocyclate) and 0.8 g of dibutyltinlaurate solution in acetone (at 10% concentration) as resetton catalyst. The reaction mixture was heated up to 90°C with stirring, and the indensation process was maintained until the isocyanate content reached 1.58 med The polyurethane prepolymer was cooled

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down to 70°C, and 30.3 g of triethylamin, wein added as neutralizing agent until a homogenous solution occurred.

The prepolymer solution was transferred over 5 minutes to a dispersion vessel equipped with a high shear cowless-type stater and containing 1265.2 g of water (at 30°C) and 0.2 g of Dehydran 1293 (anti-pam). The dispersion of the polymer was complete after about 10 minutes of stiring, and the resulting product was cooled down below 20°C. 68.1 g of 2-methyl-partanediamine dissolved in 68.1 g of water were slowly added under efficient agitation to complete the chain extension and left for 3 hours more. The aqueous dispersion of affully reacted polyurethane-urea was filtered an a  $100\mu$  sieve to deliver a stable polylirethane dispersion with a dry content of about 30.0 %, a viscosity of 100 mPa. a ph of about 8.0, a particle size of about 50 nm and a grits content below 100 mg/

Comparative example. Reference synthesis of PU 2 with standard process using 15 acetone.

isocyanate content reached 1.58 meg/g. homogenous solution occurred.

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equipped with a high shear cowless-type sitais and containing 1265.2 g of water (at S0°C) and 0.2 g of Dehydran 1293 (an foars). The dispersion of the polymer was complete after about 10 minutes of stilling, and the resulting product was cooled down below 20°C. 68.1 g of 2-methyl-pentanediamine dissolved in 68.1 g of water were slowly added under efficient agreatication to complete the chain extension and left for 3 hour more. The polyurethane dispers in was transferred to the stripping vessel, then warmed up to 50°C and the accorne was stripped out under vacuum and agitation until the free acetone level reactes a value below 0.15%. After about 1 hour of a gentle agitation, the aqueous dispers in of a fully reacted polyurethane-urea were

A double-wall glass reactor equipped with a mechanical stirrer, a thermocouple, a vapor condenser and a dropping funnel was charged with 216.2 g of acetone, 141.5 g of a polyester diol having an average molecular weight ~670 Daltons and obtained by the polycondensation of adipic acid a neopentylglycol, 42.6 g of cyclohexane dimethanol, 41.0 g of dimethylol propion acc, 383.9 g of methylene bis(cycloheny) isocyanate) and 0.8 g of dibutyltinlaurate solution in acetone (at 10% concentration) as reaction catalyst. The reaction mixtue was heated up to the boiling point of the solvent (56°C) with stirring, and the contensation process was maintained until the en, the prepolymer solution is cooled down to 50°C, and 30.3 g of triethylamine vere added as neutralizing agent until a

The prepolymer solution was transfer over 5 minutes to a dispersion vessel filtered on a 100µ sieve to deliver a stable polygrethane dispersion with a dry content

of about 30.0 %, a viscosity of 100 mpa. a pill of about 8, a particle size of about 50 nm and a grits content below 100 mg/l.

Example Reference synthesis of PU 3 via novel "coalescent-in-synthesis" process using Dowanol PGDA.

A double-wall glass reactor equipped will a mechanical stirrer, a thermocouple, a vapor condenser and a dropping funnel sas charged with 91.0 g of Dowanol PGDA, 141.5 g of a polyester diol having an a rage molecular weight ~670 Daltons and obtained by the polycondensation of a pic acid and neopentylglycol, 42.6 g of cyclohexane dimethanol, 41.0 g of dime ylol propionic acid, 383.9 g of methylene bis(cyclohexyl isocyanate) and 0.7 g of d utylindaurate solution in acetone (at 10% concentration) as reaction catalyst. The reaction mixture was heated up to 90°C with stirring, and the condensation process was maintained until the isocyanate content reached 1.86 meq/g. The polyurethane epolymer was cooled down to 70°C, and 30.3 g of triethylamine were added as new alizing agent until a homogenous solution occurred. 1390.3 g of water (at 30°C) ar 0.2 g of Dehydran 1293 (anti-foam) were added under vigorous agitation to the president properties of the phase inversion point. After about 10 minutes of stiming, the dispersion of the polymer was complete and the resulting product was cooled below 20°C. 68.1 g of 2-methyl-pentanediamine dissolved in 68.1 g of water were slowly acced under efficient agitation to complete the chain extension and left for 1 hour more witer about 3 hour of a gentle agitation, the aqueous dispersion of a fully reacted poly retbane-urea were filtered an a 100 µ sieve to deliver a stable polyurethane dispers in with a dry content of about 30 %. a viscosity of about 100 mPa.s, a pH of about 8, a particle size of about 100 nm and a grits content below 100 mg/l.

Example. In this synthesis of PU 4 with the navel "coalescent-in-synthesis" process, there is a variation of the coalescing solvent (Proglyde DMM) and the chain extender (m-xylylenediamine).

A double-wall glass reactor equipped with a mechanical stirrer, a thermocouple, a vapor condenser and a dropping firmel as charged with 104 g of Proglyde DMM, 161.7 g of a polyester diol having an average molecular weight -670 Daltons and obtained by the polycondensation of a pic acid and neopentylglycol. 48.6 g of cyclohexane dimethanol, 46.9 g of dimet vloi propionic acid. 438.8 g of methylene bis(cyclohexyl isocyanate) and 0.8 g of directly incomplete the polycondensation of a pic acid and neopentylglycol. 48.6 g of cyclohexyl isocyanate) and 0.8 g of directly incomplete the polycondensation of a pic acid and neopentylglycol. 48.6 g of cyclohexyl isocyanate) and 0.8 g of directly incomplete the polycondensation of a pic acid and neopentylglycol. 48.6 g of cyclohexyl isocyanate) and 0.8 g of directly incomplete the polycondensation of a pic acid and neopentylglycol. 48.6 g of cyclohexyl isocyanate) and 0.8 g of directly incomplete the polycondensation of a pic acid and neopentylglycol. 48.6 g of cyclohexyl isocyanate) and 0.8 g of directly incomplete the polycondensation of a pic acid and neopentylglycol. 48.6 g of cyclohexyl isocyanate) and 0.8 g of directly incomplete the polycondensation of a pic acid and neopentylglycol. 48.6 g of cyclohexyl isocyanate and 0.8 g of directly incomplete the polycondensation of a pic acid and neopentylglycol. 48.6 g of cyclohexyl isocyanate and 0.8 g of directly incomplete the polycondensation of a pic acid and neopentylglycol.

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were added under vigorous agitation t complete and the resulting product

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Example. In this synthesis of PU 5 with the new roalescent-in-synthesis" process, prepolymer molecular weight and a reduced urefnane weight contents.

obtained by the polycondensation of reaction mixture was heated up to 90 process was maintained until the iso to the polymer solution beyond the pha and a grits content below 100 mg/l

stirring, and the polycondensation pro ss was maintained until the isocyanate content reached 1.86 meq/g. The polyure mane prepolymer was cooled down to 70°C, and 35.0 g of triethylamine were added as neutralizing agent until a homogenous solution occurred. 1402.9 g of water (at 3 10) and 0.8 g of Dehydran 1293 (anti-foam) the polymer solution beyond the phase inversion point. After about 5 minutes of stirring, the dispersion of the polymer was as gooled below 20°C. 100.7 g of m xylylenediamine dissolved in 302.2 g d water were slowly added under efficient agitation to complete the chain extension and left for 3 hours of a gentle agitation. The aqueous dispersion of a fully reacted polymethane-urea was filtered on a 200 $\mu$  sieve to deliver a stable polyurethane dispersion with a dry content ca 30 %, a viscosity ca 100 mPa.s, a pH ca 8, a particle size ca linn and a grits content below 100 mg/l.

there is a reduction of the isocyamate/ drown ratio (1.6) implying and increased

A double-wall glass reactor equipped will a mechanical stirrer, a thermocouple, a vapor condenser and a dropping finne was charged with 91 g of Proglyde DMM, 258.4 g of a polyester diel having an arrage molecular weight ~670 Daltons and pic acid and neopentylelycol, 41.4 g of dimethylol propionic acid, 309.2 g of met plene bis(cyclohexyl isocyanate) and 0.7 g of dibutyltinlaurate solution in acetone (at 10% concentration) as reaction catalyst. The with stirring, and the polycondensation anate content reached 1.39 meq/g. The polyurethane prepolymer was cooled down to gooc, and 30.6 g of triethylamine were added as neutralizing agent until a home enous solution occurred. 1236.6 g of water (at 30°C) and 0.7 g of Dehydran 1293 (au foam) were added under vigorous agitation rsion point. After about 5 minutes of stirring, the dispersion of the polymer was complete and the resulting product was cooled below 20°C. 59.6 g of m-xylylen manne dissolved in 178.6 g of water were slowly added under efficient agitation to complete the chain extension and left for 3 hours of a gentle agitation. The aqueou dispersion of a fully reacted polyurethaneurea was filtered on a 200 µ sieve to del er a stable polyurethane dispersion with a dry content ca 30 %, a viscosity ca 100 Pass a pH ca 8, a particle size ca 100 nm

Example. In this synthesis of PU 6 with the novel "coalescent-in-synthesis" process, there is a stoechiometric replacement of the dimethylol propionic acid (DMPA) by the dimethylol butanoic acid (DMBA).

A double-wall glass reactor equipped with a mechanical stirrer, a thermocouple, a 5 vapor condenser and a dropping funnel was charged with 91 g of Proglyde DMM. 256.6 g of a polyester diol having an average molecular weight ~670 Daltons and obtained by the polycondensation of alipic acid and neopentylglycol, 45.4 g of dimethylol propionic acid, 307.0 g of methylene bis(cyclohexyl isocyanate) and 0.7 g of dibutyltinlaurate solution in acetone (at 10% concentration) as reaction catalyst. The 10 reaction mixture was heated up to 90% with stirring, and the polycondensation process was maintained until the isocianate content reached 1.38 meq/g. The polyurethane prepolymer was cooled down to 70°C, and 30.6 g of triethylamine were added as neutralizing agent until a homogenous solution occurred. 1245.1 g of water (at 30°C) and 0.7 g of Dehydran 1293 (an) foam) were added under vigorous agitation to the polymer solution beyond the phase inversion point. After about 5 minutes of stirring, the dispersion of the polymer vas complete and the resulting product was cooled below 20°C. 56.7 g of m-xylvlene iamine dissolved in 170.2 g of water were slowly added under efficient agitation to complete the chain extension and left for 3 hours of a gentle agitation. The aqueous dispersion of a fully reacted polyurethaneurea was filtered on a 200 sleve to deliver a stable polyurethane dispersion with a dry content ca 30 %, a viscosity ca 100 Pais a pH ca 8, a particle size ca 100 nm and a grits content below 100 mg/l.

Example. In this synthesis of PU 7 with the inevel "coalescent-in-synthesis" process. 25 there is an addition of trimethylolpropane IMP resulting in a branched prepolymer.

A double-wall glass reactor equipped with a mechanical stirrer, a thermocouple, a vapor condenser and a dropping fennel was charged with 91 g of Proglyde DMM, 228.0 g of a polyester diol having an arrage molecular weight ~670 Daltons and obtained by the polycondensation of a ipic acid and neopentylglycol, 44.6 g of dimethylol propionic acid, 6.1 g of methylolpropane, 330.3 g of methylene bis(cyclohexyl isocyanate) and 0.7 g of deputylphlaurate solution in acetone (at 10% concentration) as reaction catalyst. The reaction mixture was heated up to 90°C with stirring, and the polycondensation process was maintained until the isocyanate content reached 1.47 meq/g. The polyure rand prepolymer was cooled down to 70°C, and 33.6 g of triethylamine were added as neutralizing agent until a homogenous solution occurred. 1252.8 g of water (at Sec) and 0.7 g of Dehydran 1293 (anti-foam)

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were added under vigorous agitation, the polymer solution beyond the phase inversion point. After about 5 minutes of stirring, the dispersion of the polymer was complete and the resulting product was cooled below 20°C. 64.2 g of m-xylylenediamine dissolved in 192.5 g, waiter were slowly added under efficient agitation to complete the chain extension and left for 3 hours of a gentle agitation. The aqueous dispersion of a fully reacted popular than every as filtered on a  $200\mu$  sleve to deliver a stable polyurethane dispersion with a dry content ca 30 %, a viscosity ca 100 mPa.s, a pH ca 8, a particle size ca 10 nm and a grits content below 100 mg/1.

Example. In this synthesis of PU 8 with the novel "coalescent-in-synthesis" process, there is a neutralization of the polymer with caustic soda instead of triethylamine (TEA).

A double-wall glass reactor equipped with a mechanical stirrer, a thermocouple, a vapor condenser and a dropping finne was charged with 91 g of Proglyde DMM, 258.4 g of a polyester diol having an are erage molecular weight ~670 Daltons and obtained by the polycondensation of lipic acid and neopentylglycol, 41.4 g of dimethylol propionic acid, 309.2 g of methylene bis(cyclohexyl isocyanate) and 0.7 g of dibutyltinlaurate solution in acetone (at 10% concentration) as reaction catalyst. The reaction mixture was heated up to 90 with stirring, and the polycondensation process was maintained until the iso anaec content reached 1.39 meg/g. The polyurethane prepolymer was cooled dwn to 70°C. 12.2 g of sodium hydroxide dissolved in 420.0 g of water (at 30 g was added as neutralizing agent until a homogenous solution occurred. The remaining 840.0 g of water (at 30°C) and 0.7 g of Dehydran 1293 (anti-foam) were added valder vigorous agitation to the reactor beyond the phase inversion point. After about minutes of stirring, the dispersion of the polymer was complete and the resulting roduct was cooled below 20°C. 58.8 g of mxylylenediamine dissolved in 176 g water were slowly added under efficient agitation to complete the chain extension and left for 3 hours of a gentle agitation. The aqueous dispersion of a fully reacted perfurethane-urea was filtered on a 200 usieve to deliver a stable polyurethane dispersing with a dry content ca 30 %, a viscosity ca 100 mPa.s, a pH ca 8, a particle size ca 110 nm and a grits content below 100 mg/L

Comparative example. Reference synthesis of ACRYL 1 using standard process without AAEM.

28.6 g of an aqueous solution of sodit in nonylphenylpoly(exyethylene) sulfate with n=10 (solids content of 34wt%) at 1 28.6 g of an aqueous solution of

Empfansszeit 14.Juli 14:27:

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nonylphenoxypoly(oxyethylene) with next (solids content of 70wt%) and 5.0 g of the potassium salt of 3-sulfopropyl methacin ate were introduced with stirring in a tank containing 290.0 g of demineralized water. Then, 575.0 g of methyl methacrylate, 410.0 g of 2-ethylhexyl acrylate and 15.0 g of acrylic acid were added thereto with strong stirring, and resulting in the formation of a preemulsion. 2.4 g of ammonium persulfate were added with stirring to a reactor containing 4.3 g of the abovementioned aqueous solution of nonylic enylpoly(oxyethylene)sulfate in 720.0 g of demineralized water and heated up to 5°C. The pre-emulsion prepared above was then added into the resulting mixture over a period of 2.5 hours. The reactor was maintained at 80°C. for 2 hours to come ste the reaction and then allowed to cool to room temperature. 10.0 g of a 25% (w/v) aqueous solution of ammonia were added slowly thereto. The resulting latex had a ry content of 48.0%, a viscosity of 315 mPa s, a pH of 8.5, an average particle size of 134 nm, a free monomer content of below 0.01wt%, a grits content below 50 mg/s and a minimal film forming temperature of about 17°C. This vinyl polymer had no across the polymer functional groups.

Example. Reference synthesis of ACRYL pusing standard process with AAEM.

28.6 g of an aqueous solution of sodium nonylphenylpoly(oxyethylene)sulfate with n=10 (solids content of 34wt%) and 28.6 g of an aqueous solution of nonylphenoxypoly(oxyethylene) with n=86 (solids content of 70wt%) and 5.0 g of the potassium salt of 3-sulfopropyl methacitate were introduced with stirring in a tank containing 290.0 g of demineralized water. Then, 550.0 g of methyl methacrylate. 385.0 g of 2-ethylhexyl acrylate, 500 g acefoacetoxyethyl methacrylate and 15.0 g of acrylic acid were added thereto with rong stirring, and resulting in the formation of a preemulsion. 2.4 g of ammonism persulfate were added with stirring to a reactor containing 4.3 the ve-mentioned aqueous nonylphenylpoly(oxyethylene)sulfate in 720.0 g of demineralized water and heated up to 80°C. The pre-emulsion prepared about was then added into the resulting mixture over a period of 2.5 hours. The reaction was maintained at 80°C. for 2 hours to complete the reaction and then allewed cool to room temperature. 10.0 g of a 25% (w/w) aqueous solution of ammonia with added slowly thereto. The resulting latex had a dry content of 48.6%, a viscosity 232 mPa s, a pH of 6.0, an average particle size of 133 nm, a free monomer coment of below 0.01wt% (controlled by gas chromatography), a grits content belo 50 mg/l and a minimal film forming temperature of about 20°C.

Example. Synthesis of ACRYL Susting cont & shell process with AAEM.

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51.3 g of an aqueous solution of sodium nonylphenylpoly(oxyethylene) sulfate with n=10 (solids content of 30wton and 148 g of an aqueous solution of 72.7 g of water. The reaction mixsur average particle size of 110 nm, a free moreomer content of below 0.01wt% (controlled by gas chromatography), a grite content clow 50 mg/l and a minimal film forming

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148 g of an aqueous solution of nonylphenoxypoly(oxyethylene) with n=36 (solids content of 80wt%) were introduced with stirring in the reactor containing 2073.8 g of demineralized water. 444.6 g of a seed latex made from the radical polymerisation of methyl methacrylate (3), styrene (54.2), butyl acrylate (23.6), 2-etisylhes lacrylate (17.7) and methacrilic acid and having a dry content of 37%, a pH of 7.5 and an almost homodisperse particle size of 57 nm. The temperature is raised to 80% and 2.3 g of sodium persulfate is added to the reactor. A mixture of 1062.3 g of statene and 38.5 g of 2-ethylhexyl acrylate is added during 1h30' with a parallel addition of 3.4 g of sodium persulfate dissolved in ure in allowed to react for a further 30' under agitation. Then, a mixture of 71.9 g of 2 hylhexyl acrylate. 575.6 g of butyl acrylate. 43.2 g of methacrylic acid and 43.2 g of agetoacetoxyethyl methacrylate were added to the reactor in a period of 1h. The reactor was maintained at 80°C for 1 hour to complete the reaction and then allowed in cool to room temperature. 25.0 g of a 25% red frool to room temperature. 25.0 g of a 25% (w/w) aqueous solution of ammonia were added slowly to neutralize the latex. The resulting product had a dry content of 4412, a viscosity of 100 mPa s. a pH of 7.0, an

Example. Synthesis of ACRYL 4 with a standard modified process and with a decrease of the glass transition temperature ag by the reduction of the methyl methacrylate / 2-ethylhexyl acrylate ratio; the acetoace oxy- and carboxylic acid functionality is maintained.

38.3 g of an aqueous solution of dodecyl mydibenzene disulfonate, sodium salt (solids content of 45wt%) is introduced with stirring in a tank containing 361 g of demineralized water. Then. 517.5 g of reschyl inethacrylate, 557.8 g of 2-ethylhexyl acrylate, 57.5 g of acetoacetoxyethin methacrylate and 17.3 g of acrylic acid were added consecutively with a strong aggration to form a stable monomer pre-emulsion. A reactor is charged with 828 g of dem neralized water and 3.8 g of an aqueous solution of dodecyl oxydibenzene disulfornie. sodium salt (solids content of 45wt%) under an efficient agitation. The reaction vessel is heated up to 80°C and 2.9 g of potassium persulfate is added as the initiator. The pre-emulsion prepared above is then loaded into the reactor over a period 2 hours, and the reactor is maintained at

80°C for 2 hours to complete the reaction. The reactor then allowed to cool to room

temperature and 3.8 g of a 25% (w/w) as leous solution of ammonia is slowly added a minimal film forming temperature belog 5°C.

for neutralization. The resulting latex had a dry content of 49%, a viscosity ca 200 mPa.s, a pH ca 6.0, an average particle size ca 130 nm, a free monomer content of below 0.01wt% (controlled by gas chromatography), a grits content below 50 mg/l and

Example. Synthesis of ACRYL 5 with a standard modified process and with a neutralization of the polymer with causting oda instead of ammonia.

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76.7 g of an aqueous solution of dodecylingydibenzene disulfonate, sodium salt (solids 10

content of 45wt%) is introduced with stirring in a tank containing 693 g of demineralized water. Then, 1265.0 g of methyl methacrylate, 885.5 g of 2-ethylhexyl acrylate, 115.0 g of acetoacetoxyethyl methacrylate and 34.5 g of acrylic acid were added consecutively with a strong agitation to form a stable monomer pre-emulsion.

A reactor is charged with 1656 g of demineralized water and 7.7 g of an aqueous solution of dodecyl oxydibenzene disulferate; sodium salt (solids content of 45wt%) and a minimal film forming temperature 20°C.

under an efficient agitation. The reaction vessel is heated up to 80°C and 5.5 g of potassium persulfate is added as the initiator. The pre-emulsion prepared above is then loaded into the reactor over a period of 2.5 hours, and the reactor is maintained at 80°C for 2 hours to complete the reaction. The reactor then allowed to cool to room temperature and 115 g of a 5% (w/w) agreeous solution of sodium hydroxide is slowly added for neutralization. The resulting per had a dry content of 48%, a viscosity ca 200 mPa.s, a pH ca 6.0, an average particle size ca 130 nm, a free monomer content of below 0.01wt% (controlled by gas chritinatography), a grits content below 50 mg/l

Examples 1 to 6

The polyurethane dispersions (PU 1-3) have been assessed alone as references 1-5 and combined with acrylic latexes in bands 1-6 as outlined in the table 1, 2 and according to the procedure described above.

	POLYURETHANE POLYMER	ACRYLIC POLYMER	COALESCENT ADDED after synthesis
Reference 1	· PU 1		
	(NMP) ;	L ii	
Reference 2	PU 2 .	_	
	(ACETONE)		

Empfangszeit 14.Juli 14:27

Reference 3	PU 2 : (ACETONE)		•	8% Dowanoi EB/DPM 1:3
Reference 4	PU 3 (PGDA coalescent)	<u> </u>	-	25/5/11:5
Reference 5	PU 3 (PGDA coalescent)		-	8% Dowanol EB/DPM 1:3

Table 1: polyurethane references (PU 1-3)

			. 15	
(50:50 dry blend)	POLYURETHANE POLYMER :		ACRYLIC	COALESCENT ADDED after synthesis
Blend 1	PU 1     (NMP)		(NO AAEM)	-
Blend 2	PU 1 (NMP)		ACRYL 2	-
Blend 3	PU 1 (NMP)	. :	ACRYL 3	
Blend 4	PU 2 . (ACETONE)		ACRYL 2	-
Blend 5	FU 2 (ACETONE)			4% Dowanol PGDA
Blend 6	PU 3 (PGDA coalescent)		ACRYL 2 (AAEM)	-

Table 2: polyurethane : acrylic hybrid control positions from PU 1-3

The different polyurethane dispersions were applied on glass and assessed for König hardness as disclosed in table 3.

		<del></del>		齈				
	Ratio %	1h			4h.	24h	4 days	7 days
	1h/7d				]· : [[			, days
Reference 1	6	10	: 1		22	69	138	151
Mfft 15°C			[[] :		1 11		136	151
Reference 2		NO FILM	FOE	ij,	ATION A	T POOM TEN	PERATURE	
Mft>70°C						I MOOM IEM	PERATURE	
Reference 3	41	52	:		61,	87	140	107
			1				1=0	127

Empfangszeit 14.Juli 14:27

				1 !!!			
Reference 4 Mfft 63°C		NO FILM	FOR	ATION A	T ROOM TEM	PERATURE	•
Reference 5	90	94		90	93	98	105

Table 3: König kinetics of drying from PU 1-3

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The different blends were applied on the several test substrates and assessed for performance as disclosed in tables 4 and p

			KU	1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	_		
	Ratio % lh/7d	1h		4h	24h	4 days	7 days
Blend 1 (Mfff: 6°C)	16	15		25	51	77	92
Blend 2 (Mfft: 6°C)	15	13		22	52	66	84
Blend 3 (Mfft: 7°C)	8	10		36	91	121	133
Blend 4 (Mfft: 30°)	49	36	H	43	54	69	74
Blend 5 (Mfft: 0°C)	61	46		51.	59	70	. 75
Blend 6 (Mfft: 18°)	. 95	104		99	105	107	109

Table 4: König kinetics of drying from Plili-3

	··-				
	Clarity	Stain	Ethanol	Scratch	Adhesion
•	(1-5, best)	resistance	tesistance	resistance	dry - wet
		(1-5, best)	(d.rubs)	(1-5, best)	(1-5, best)
Blend I	4.	3.7	20	5?	4-4
Blend 2	· 4	4.3	>100	5	4-5
·Blend 3	2	1.8	>100	, 4	5-5
Blend 4	4	3.5	>100	2	4-3
Blend 5	4:	4.0	>100	5	4-5
Blend 6	3.5	4.0	>100	5	4-5

Table 5: Properties & performance on ward for coatings from PU 1-3

Empfanøszeit 14.Juli 14:27.

## Examples 7 to 12

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The polymer dispersions have been contained in examples 7-12 as outlined in the table 4 and according to the procedure rescribed above. They further exemplify the scope of the invention.

			3 1	44:	
(50:50 dry blend)	POLYURETHANE POLYMER			ACRYLIC POLYMER	COALESCENT ADDED after synthesis
Blend 7	PU 4 (Proglyde DMM coalescent)	Section 200		ACRYL 2	4% Dowanol EB/DPM
Blend 8	PU 5 (Proglyde DMM coalescent)	THE PERSON NAMED IN		ACRYL 2	1% Proglyde DMM
Blend 9	PU 6 (Proglyde DMM coalescent)			ACRYL 2	1% Proglyde DMM
Blend 10	PU 7 (Proglyde DMM coalescent)			ACRYL 2	1% Proglyde DMM
Blend 11	PU 8 (Proglyde DIMM coalescent)	The second second		ACRYL 5	1% Proglyde DMM
Blend 12	PU 5 (Proglyde DMM coalescent)	the second second		ACRYL 4	

Table 6: polyurethane : acrylic hybrid compositions from PU 4-8

The different blends were applied on the sest substrates and assessed for performance as disclosed in tables 5 and 6.

			_ I H (				
	Ratio %	lh.		4h	24h	4 days	7 days
•	1h/7d			<u> </u>	1		- 4.32
Blend 7	85	90		94	97	101	106
(Mfft:17°C)				117	' "'	1 .02	108
Blend 8	69	59	!	66	66	84	. 96
(Mfft: 9°C)							· 86
Blend 9	68	61		64	71	75	90
(Mfft: 2°C)					• • •	, ,	90

Empfansszeit 14. Juli 14:27

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Blend 10	75 ·	75	78	T		
(Mfft: 8°)			78	84	88	100
Blend 11	78	87	88	300		
(Mfft: 8°C)				100	-	112
Blend 12	64	39	44	50		
(Mfft: 0°C)			44	52	61	61
			:			

Table 7: König kinetics of drying from Pu

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		·		[·		•	
	Clarity	Stain	Aceto		Scratch	Adhesion	RT film
	(1-5, best)	resistance	resista		resistance	dry - wet	formation
		(1-5, best)	d.ru	bန္က)	(1-5, best)	(I-5, best)	(1-5, best)
Blend 7	2,25	3.5	85	4:	4	1-1.5	
Blend 8	3.75	3.8	50	11			4.5
Blend 9	3.75	3.58	Rilla	<del></del>	4	5-5	4.5
Blend 10					2	4-5	4
	3.5	4.25	90	i	3	3.5-4.5	3.5
Blend 11	4.0	3.35	60	1	2.5		
Blend 12	2.25		•.03011		2.0	3-4.5	3
Table 8: Pro	perties & perfo			11 1		-	4.5

Table 8: Properties & performance on would for chatings from PU 4-8

One can see from the examples that the polyure hane component containing the NMP of the chemical resistance can be achieved lower chemical resistance.

(blends 1-3) failed to give a satisfactory fast hardness development. The improvement d with a self-crosslinking system (blend 2) while the improvement of the final hardness can be obtained with a core & shell morphology (blend 3) without detrimental effect on the film formation, but despite the

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The total suppression of the NMP in the polyurethane component improves the last hardness (blend 4) which is still the case when a coalescing solvent is added to the product to descrease the MFFT (blend ). But the best contribution to the fast hardness is obtained with the comparative coalescent-in-synthesis process (blend 6) that does not significantly depress the in formation and the performance of the coating on wood.

The composition of both the polyurethane and the acrylic components can be modified in large proportions in order to optimize the overall performance of the products and dipropyleneglycol dimethylether (Proglyc diamine (MXDA) is very efficient to rec favorable to obtain a good transparen propionic acid (blend 8) or dimethyle resistance through polymer branching (thend 10). It is possible to obtain an aminefree composition by the replacement of the amine with sodium hydroxide in the polyurethane dispersion and in the acrysic latex (blend 11); the favorable effect on occupational health is balanced by a lower resistance of the film. The use of an acrylic with another backbone composition provides endless combinations with the polyurethane dispersion; the reduction of the acrylic polymer is for instance favorable for the film formation without increasing the coalescing solvent contents, but the transparency and the solvent resistance is somewhat reduced accordingly (blend 12).

to influence positively the combination of antagonistic properties, like low film formation temperature and high hadness. It is shown that the use of ness. It is shown that the use of DWW) in combination with m-xylylene d a fast hardness with a stable polymer dispersion (blend 7). The decrease of the isocyanate / hydroxyl functionality is and adhesion, either when dimethylol buignoic acid (blend 9) is used. The incorporation of trimethylolpropane provides a favorable increase of the hardness and

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### Claims

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- 1. A composition comprising:
  - (A) a dispersion of a crosslinkable polygrethane polymer in aqueous medium, and obtainable from a polygrethane prepolymer which is the reaction product of:
- (1) at least one polyisocyanate, and
- (ii) at least one organic compound containing at least two reactive groups which can react with isocyanates, and
- at least one compound which is capable to react with an isocyanate group and which contains additional functional groups which are susceptible either to dispersion in water and/or to a cross-linking reaction
  - (iv) in an oxygenated solvent (iv) selected from the coalescing agents having a higher boiling point of from 150 to 250°C, under 760 mm Hg and being chemically inert towards isocyamates, during the manufacture of the polyurethane;
    - the so obtained polyurethane prepolymer being further neutralised and dispersed in water, then reacted with a chain extender capping agent (v) having or not remaining functional groups after the capping:
  - (B) a crosslinker, if the polyurethane is not self-crosslinkable
  - 2. A composition according to claim 1, wherein said oxygenated coalescing solvent (iv) is selected from the fully reacted alient or aryl esters of aromatic, aliphatic or cycloaliphatic polyglycols, the fully reacted alkyl or aryl esters of aromatic, aliphatic or cycloaliphatic polyglycols, the fully reacted alkyl or aryl ethers of aromatic, aliphatic or cycloaliphatic polyglycols, the fully reacted mixed alkyl and aryl esters and ethers of mixed aromatic, aliphatic or cycloaliphatic polyglycol-carboxylates, the neat, the alkyl and aryl substituted cyclic carbonates, the neat, the alkyl and aryl substituted cyclic esters, the neat, the alkyl and aryl substituted cyclic anhydrides.
- 3 A composition according to claim 2, wherein said oxygenated coalescing solvent (iv) is selected from dimethyl esters or discountyl esters of adipic, glutaric, succinic or phtalic acids, the ethyl-3-ethoxypropionate, the 2.2,4-trimethyl-1.3-pentanedioldisobutirate, the ethylene carbonate, the propylene carbonate, the propylene carbonate, the propyleneglycol diacetate and the dipropylene glycol dimethyl ether, alone or in admixture

- A composition according to any of claims 1 to 4, wherein the organic polyisocyanate is an aliphatic, cycloal phatic, aromatic or heterocyclic polyisocyanate.
- 6 A composition according to any of claims 1 to 5 wherein the organic compound (ii) containing at least two isocyanate-reactive groups is selected from the group consisting of polyester polyols, polyether polyols, polycarbonate polyols, polyacetal polyols, polyesteramide polyols, polyacetal polyols, polyesteramide polyols, polyacetal alone or in admixture.

A composition according to any of claims 1 to 6, wherein the isocyanate-reactive compound (iii) containing the anionic salt functional groups, or acid groups which may be subsequently converted to sumb anionic salt groups, is represented by (a) a hydroxycarboxylic acid of general formula (HO)<sub>x</sub>R(COOH)<sub>y</sub>, wherein R represents a straight or branched chain hydrocarbon radical having 1 to 12 carbon atoms, and x and y are integers from 1 to 3, or (b) a suifonated polyester obtained by the reaction of a sulfonated dicarboxylic acid with one or more polyhydric alcohols, or by the reaction of a sulfonated diol with one or more polycarboxylic acids.

- A composition according to any of claims 1 to 7, wherein the capping agent (v) is water or an aliphatic, alicyclic, aromatic or heterocyclic primary or secondary polyamine having up to 80 carbon atoms which can bear a further functional goup such as in gamma-aminopropyltrime hoxysilane, gamma-aminopropyltriethoxysilane, N-beta-aminoethyl-gamma-aminopropyltrimethoxysilane, bis-(gamma-trimethoxysilylpropyl) amine and N-beta-(aminoethyl)-gamma-aminopropylmethyldimethoxysilane.
- 9 A composition according to any of claims I to 8, wherein said at least one vinyl polymer is the product formed by the free-radical addition polymerization of at least one monoethylenically unsaturated monomer with at least one other ethylenically unsaturated monomer containing a functional group such as an acetoacetoxyalkyl ester group, carboxylic and sulfonic groups, isocyanates, hydroxy, amine, acrylic, allylic, vinyl alkenyl, alkinyl, halogen, epoxy, aziridine, aldehyde, ketone, anhydride, carbonate, stane, carbodiimide, ureidoalkyl, N-methylolamine, N-methylolamide N-alkoxy-methyl-amine, N-alkoxy-methyl-amide, and capable to provide a crosslinking reaction with the polyurethane component.

10 A composition according to any of claim 1 to 9, wherein the monoethylenically unsaturated monomer containing an acetoacetoxyalkyl ester group is a compound having the formula

R-O-CO-CH2-CO-CH3

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wherein R represents a

CH<sub>2</sub>=CR'-COO-R"- group or a CH<sub>2</sub>=CR'-R"- group in which

R' is a hydrogen atom or a methyl radical and

R" is an alkylene radical maving 1 to 12 carbon atoms.

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- A composition according to any of claims to 10, wherein the monoethylenically unsaturated monomer containing an acctoacetoxyalkyl ester group is acctoacetoxyethyl (meth)acrylate.
- 15 12 A composition according to any of claims into 11, wherein the monoethylenically unsaturated monomer containing an accetoacetoxyalkyl ester group is present in an amount of from about 1 to about 80 % by weight of the vinyl polymer.
- A composition according to any of claims 1 to 12, wherein the monoethylenically unsaturated monomer containing an acetoacetoxyalkyl ester group is present in an amount of from about 5 to 20 % by weight of the vinyl polymer.
- A composition according to any of claims 1 to 13, wherein the weight ratio of said at least one polyurethane polymer to said at least one vinyl polymer is within the range of from 10:0 to 1:10.
  - A composition according to any of claims 1 to 14, wherein the weight ratio of said at least one polyurethane polymer to said at least one vinyl polymer is within the range of from 1:2 to 2:1.

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16. A process for the preparation of an aqueous selfcrosslinkable resin composition according to any of claims 1 to 15, which comprises homogenously mixing together at room temperature an aqueous dispersion of at least one polyurethane polymer and an aqueous dispersion of at least one vinyl polymer

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17. A process for the preparation of an adjustous selfcrosslinkable resin composition according to any of claims 1 to 15, which comprises subjecting the monomers of the vinyl polymer having functional groups to radical polymerisation in the

Empfangszeit 14. Juli 14:27

presence of an aqueous polyurethane polymer having anionic salt functional groups or subjecting an isocyanate-terminated polyurethane prepolymer having anionic salt functional groups to chain-extension with a capping agent in the presence of an aqueous dispersion of a viny polymer having functional groups.

18. A protective or adhesive coating obtained with an aqueous composition according to any of claims 1 to 17.

### **Abstract**

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The present invention concerns a composition and a process for making crosslinkable polyurethane dispersions to be used alone of as an hybrid polymer dispersion in water, said hybrid polymer dispersion being constituted by the functionalized polyurethane polymer (A) and a functionalised vinylic polymer (B) either as individual particles or as composite particles

A isocyanate-terminated or hydroxy-terminated polyurethane prepolymer is first prepared in a high boiling point oxygenated coalescing solvent having no functional group reactive with isocyanates and trat will remain after water dispersion. The composition is thereafter neutralized, dispersed in water and chain extended with a diamine.

The polymer dispersions according to the invention exhibit a low film forming temperature associated with a very fast hardness development and a good resistance to water, solvents and stains.

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